CH-420 Understanding advanced molecular simulation

Smit Berend				
Cursus	Sem.	Туре	Language of	English
Chimiste	MA2, MA4	Opt.	teaching	English
Computational science and Engineering	nal science and Engineering MA2, MA4 Opt. Credits Session	4 Summer		
			Semester	Spring
			Exam	During the semester
			Workload	120h
			Weeks	14
			Hours	3 weekly
			Courses	2 weekly
			Project	1 weekly
			Number of positions	

Summary

This course introduces advanced molecular simulation techniques such as Monte Carlo and Molecular dynamics in different ensembles, free energy calculations, rare events, Configurational-bias Monte Carlo etc.

Content

The course will be given in five blocks, each covers a topic and ends with an assignment to carry out some simulations.

- 1. [Weeks 1-3] Computational Carpentry and Statistical Thermodynamics
- 2. [Weeks 4-5] Basic Monte Carlo and Molecular Dynamics
- 3. [Weeks 6-9] Monte Carlo and Molecular dynamics in different ensembles
- 4. [Week 7] semester break
- 5. [Weeks 10-13] Free energy calculations and rare events
- 6. [Weeks 14-15] Special topics

Keywords

molecular simulation, molecular dynamics, Monte Carlo simulation

Learning Prerequisites

Recommended courses

knowledge on statistical thermodynamics, some basic programming skills

Assessment methods

The mark will be a: assignments given during the course (60%) and reports + oral presentation of 2 blocks (40%)

Resources

Bibliography

D. Frenkel and B. Smit, *Understanding Molecular Simulations: from Algorithms to Applications*, 2nd ed. (Academic Press, San Diego, 2002)

Ressources en bibliothèque

• Understanding molecular simulation : from algorithms to applications / Frenkel

